

A separate lowstand lake at the northern edge of Lake Tanganyika? Evidence from phylogeographic patterns in the cichlid genus *Tropheus*

Christian Sturmbauer  · Christine Börger ·
Maarten Van Steenberge · Stephan Koblmüller

Received: 24 February 2016 / Revised: 8 July 2016 / Accepted: 31 July 2016
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Abstract In Lake Tanganyika, lake level fluctuations were shown to have had a major impact on the evolution of littoral species. Many species are subdivided into arrays of populations, geographical races and sister species, each colonizing a particular section of the shore. Their often limited dispersal abilities promoted geographic isolation and, on the long run, allopatric speciation. With more than 120 distinct populations, the genus *Tropheus* represents the most spectacular and best-studied example of this phenomenon. The present study aims at the fine-scale reconstruction of the spread of two mitochondrial *Tropheus*-lineages in the very north of the lake, where two species, *T. sp. 'black'* and *T. brichardi*, occur. Using mtDNA sequences and AFLP-data, we ana-

lyzed samples from 21 localities and found a highly complex conglomerate of introgressed populations formed by the repeated contact of two lineages. Our data suggest repeated cross-lake dispersal of *T. sp. 'black'* haplotypes along the ridge between the West and East Ubwari Fault, supporting an additional persisting lowstand-lake in the Bujumbura subbasin at the very north of the lake and highlighting once more the impact of lake level fluctuations on the genetic structure and evolution of stenotopic rock-dwelling cichlid species.

Keywords mtDNA sequences · Control region · AFLP · Secondary admixis · Hybridization · Lake level fluctuations

Electronic supplementary material The online version of this article (doi:[10.1007/s10750-016-2939-8](https://doi.org/10.1007/s10750-016-2939-8)) contains supplementary material, which is available to authorized users.

Guest editors: S. Koblmüller, R. C. Albertson, M. J. Genner, K. M. Sefc & T. Takahashi / Advances in Cichlid Research II: Behavior, Ecology and Evolutionary Biology

C. Sturmbauer (✉) · C. Börger · M. Van Steenberge ·
S. Koblmüller
Department of Zoology, University of Graz,
Universitätsplatz 2, 8010 Graz, Austria
e-mail: christian.sturmbauer@uni-graz.at

M. Van Steenberge
Department of Ichthyology, Royal Museum for Central
Africa, Leuvensesteenweg 13, 3080 Tervuren, Belgium

Introduction

Genetic and phenotypic divergence among populations forms the basis of speciation events. However,

M. Van Steenberge
Laboratory of Biodiversity and Evolutionary Genomics,
University of Leuven, Charles Deberiotstraat 32,
3000 Leuven, Belgium

S. Koblmüller
Institute of Vertebrate Biology, Academy of Sciences of
the Czech Republic, Květná 8, 603 65 Brno, Czech
Republic

such pathways rarely proceed in a linear fashion by divergence alone, as repeated intermezzos of gene flow perturbate gene pools and set the stage for selection, drift, and speciation (King & Lawson, 1995; Rossiter, 1995; Pinho & Hey, 2010; Sturmbauer, 2011). On the long run, secondary admixis can either render population distinctness or produce novel and distinct hybrid entities, which can evolve to novel species (Seehausen, 2004; Nolte & Tautz, 2010).

Following the spatial segregation into rock- and sand-habitats along the shores of Lake Tanganyika, many stenotopic littoral species are subdivided into distinct populations that vary mainly in coloration. The genus *Tropheus* represents perhaps the best example for this phenomenon. *Tropheus* is abundant in the upper littoral zone in all types of rocky habitats, where it feeds on epilithic algae and takes shelter from predators. Sandy or muddy shores and river estuaries are strictly avoided, resulting in about 120 distinct local variants, some of which live in sympatry (Schupke, 2003; Konings, 2013). The six described species (Poll, 1986) are currently under revision (Van Steenberge, 2014).

A series of previous studies on the evolution and colonization history of this genus set out to reconstruct the origin and spread of this highly specialized rock-dwelling species-complex (Sturmbauer & Meyer, 1992; Sturmbauer et al., 1997; Rüber et al., 1999; Baric et al., 2003; Sturmbauer et al., 2005; Egger et al., 2007; Sefc et al., 2007, 2016; Koblmüller et al., 2011; Nevado et al., 2013). These studies suggested mitochondrial introgression, sometimes on a small scale between presently allopatric populations, sometimes on a large-scale producing true hybrid populations, at various time points in the evolutionary history of the genus *Tropheus*. Most of these events of population displacement were triggered by lake level fluctuations, which extended to a few hundred meters below present level (Lezzar et al., 1996; Cohen et al., 1997; Scholz et al., 2003; McGlue et al., 2008). While the effect is widely seen in population genetic studies on various Lake Tanganyika cichlid species (Verheyen et al., 1996; Rüber et al., 1999, 2001; Duftner et al., 2006; Koblmüller et al., 2007, 2009, 2011; Sefc et al., 2007; Nevado et al., 2013; Van Steenberge et al., 2015), studies specifically testing for the impact of Pleistocene water level fluctuations are scarce (Koblmüller et al., 2011; Sefc et al., 2016; Winkelmann et al., 2016).

In the present study, we aim at the fine-scale reconstruction of the origin and spread of the—in terms of colors—highly divergent *Tropheus* populations in the very north of Lake Tanganyika. In this part of the Lake, three species of *Tropheus* are found. Besides the basal species *T. duboisi* Marlier 1959, these are *T. brichardi* Nelissen & Thys van den Audenaerde 1975 and the hitherto undescribed species *T. sp. 'black'* (Konings, 2013). *Tropheus duboisi* co-occurs with *T. sp. 'black'* at Bemba, and with *T. brichardi* at Mwamugongo. At the eastern shore of the Ubwari Peninsula between Cape Caramba and Muzimu two *Tropheus* live in sympatry, one currently assigned to *T. sp. 'black'* named Caramba, and the second tentatively assigned to *T. cf. brichardi* named Ubwari-green in the aquarium trade (Schupke, 2003; Konings, 2013). We analyzed samples from 21 populations of *T. brichardi* and *T. sp. 'black'* and found a highly complex conglomerate of populations, formed by the repeated contact of two ancient mtDNA lineages, and relate the findings to patterns in nuclear DNA markers using new AFLP data.

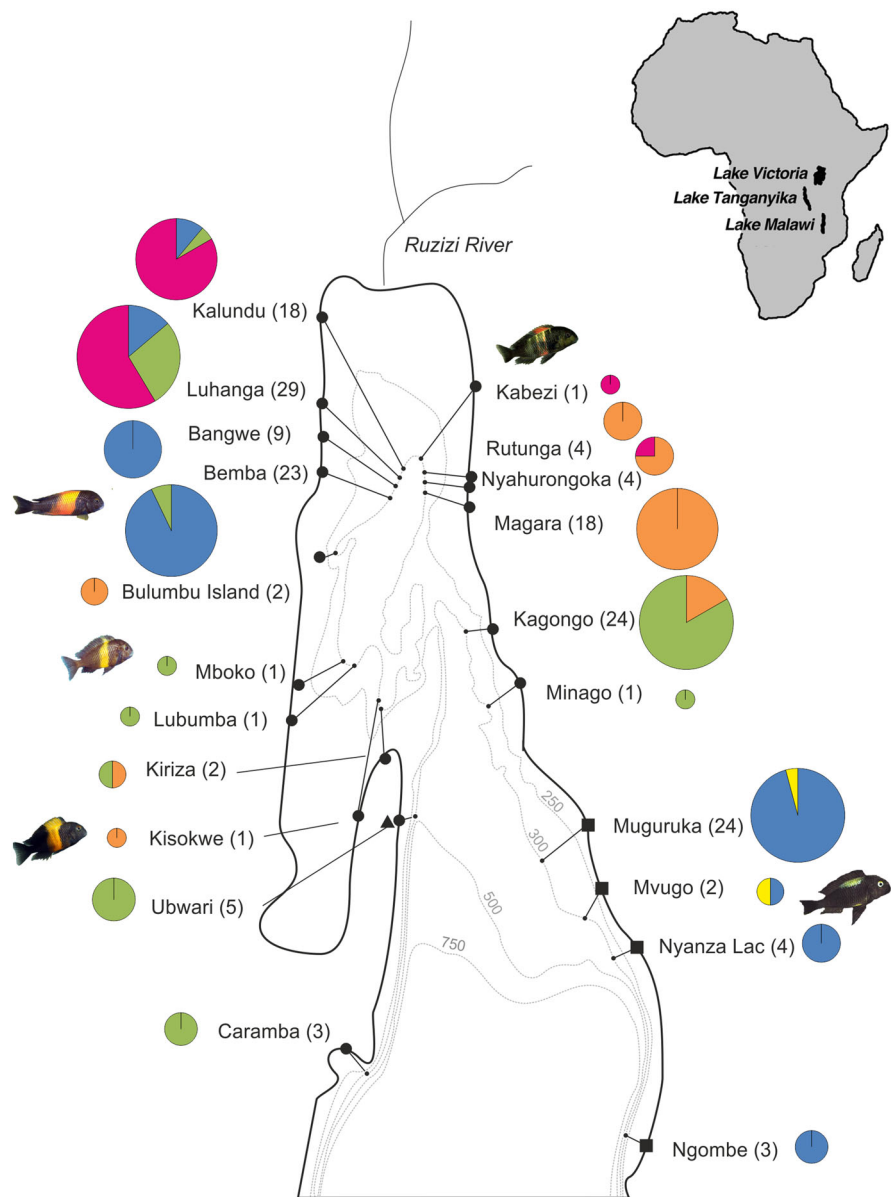
Materials and methods

Samples and molecular techniques

This study is based on 179 sequences of the most variable part of the mitochondrial control regions and AFLP profiles of 30 individuals. Fin clips of *Tropheus brichardi* and *T. sp. 'black'* were collected from 21 localities in the northernmost section of Lake Tanganyika (Fig. 1) during several expeditions between 1991 and 2013 (Van Steenberge, 2011), or obtained via the aquarium trade, and preserved in >96% ethanol. Mitochondrial control region sequences for 27 of these samples have been published previously (Sturmbauer & Meyer, 1992; Baric et al., 2003; Sturmbauer et al., 2005; Egger et al., 2007), and 152 new samples were sequenced in this study (Table 1). Whole genomic DNA was extracted following a rapid Chelex protocol (Richlen & Barber, 2005) or using the Macherey–Nagel Nucleospin extraction kit, following the manufacturer's instructions.

For all 152 new samples, the most variable part of the mitochondrial control region was amplified and sequenced following Koblmüller et al. (2011) and Duftner et al. (2005), respectively. The primers used for PCR amplification and chain termination sequencing

Fig. 1 Map of northern Lake Tanganyika with sampling sites. Numbers in parentheses refer to sample size; pie charts indicate assignment to major haplogroups (Fig. 2), with size of the pie chart proportional to the number of samples. *Square symbols* marking sampling sites indicate the occurrence of *T. brichardi* instead of *T. sp. 'black'* (circle symbols). The *triangle symbol* denotes the occurrence of *T. cf. brichardi* named Ubwari-green in the aquarium trade



were L-Pro-F (Meyer et al., 1994) and TDK-D (Lee et al., 1995). DNA fragments were purified with SephadexTM G-50 (GE Healthcare) and visualized on an ABI 3130xl automated sequencer (Applied Biosystems). Sequences were aligned using ClustalW in the computer program MEGA V6.0 (Tamura et al., 2013) and the resulting alignment was controlled by eye to check for obvious alignment errors. The final alignment length was 352 bp. The new DNA-sequences have been deposited in Genbank. Sample information and all accession numbers are listed in Table 1.

AFLP genotyping of 30 individuals (ten primer combinations for selective amplification: *EcoRI*-ACA/*MseI*-CAA, *EcoRI*-ACT/*MseI*-CAG, *EcoRI*-ACC/*MseI*-CAC, *EcoRI*-ACA/*MseI*-CAG, *EcoRI*-ACA/*MseI*-CAC, *EcoRI*-ACA/*MseI*-CAT, *EcoRI*-ACT/*MseI*-CAT, *EcoRI*-ACT/*MseI*-CAA, *EcoRI*-ACT/*MseI*-CAC, *EcoRI*-ACC/*MseI*-CAA) followed the protocol described in Egger et al. (2007). Selective PCR products were sized against an internal standard (GeneScan-500 ROX, Applied Biosystems) on an ABI 3130xl automated sequencer (Applied Biosystems). The AFLP data are part

Table 1 Characterization of the individuals studied, with information concerning sampling locations and their GPS coordinates, extraction number, GenBank accession number, species and haplotypes to which individuals were assigned as well as the sample identification

Location	Latitude (S)	Longitude (E)	Extraction no.	Accession no.	Species	Haplotype	Sample ID
Bangwe	−3,7919	29,1627	14420	KX513704	<i>T. sp.</i> ‘black’	Ht_01	KS23H4
			14423	KX513701	<i>T. sp.</i> ‘black’	Ht_02	KS23H4
			14424	KX513700	<i>T. sp.</i> ‘black’	Ht_02	KS23H5
			14454	KX513671	<i>T. sp.</i> ‘black’	Ht_02	T14/03/A4
			14455	KX513670	<i>T. sp.</i> ‘black’	Ht_02	T14/03/A5
			14419	KX513705	<i>T. sp.</i> ‘black’	Ht_03	KS23H4
			14421	KX513703	<i>T. sp.</i> ‘black’	Ht_03	KS23H4
			14422	KX513702	<i>T. sp.</i> ‘black’	Ht_03	KS23H4
			14426	KX513699	<i>T. sp.</i> ‘black’	Ht_03	KS23H5
Bemba	−3,7919	29,1627		Z12099	<i>T. sp.</i> ‘black’	Ht_01	
				Z12091	<i>T. sp.</i> ‘black’	Ht_01	
				Z12100	<i>T. sp.</i> ‘black’	Ht_01	
				Z12098	<i>T. sp.</i> ‘black’	Ht_01	
				Z72097	<i>T. sp.</i> ‘black’	Ht_01	
				Z12095	<i>T. sp.</i> ‘black’	Ht_01	
				Z12093	<i>T. sp.</i> ‘black’	Ht_01	
				Z12092	<i>T. sp.</i> ‘black’	Ht_01	
			14451	KX513674	<i>T. sp.</i> ‘black’	Ht_01	T14/03/A1
			14388	KX513736	<i>T. sp.</i> ‘black’	Ht_01	KS22J6
			14389	KX513735	<i>T. sp.</i> ‘black’	Ht_01	KS22J6
			14393	KX513731	<i>T. sp.</i> ‘black’	Ht_01	KS23A10
			14394	KX513730	<i>T. sp.</i> ‘black’	Ht_01	KS23A10
			14390	KX513734	<i>T. sp.</i> ‘black’	Ht_02	KS22J6
			14391	KX513733	<i>T. sp.</i> ‘black’	Ht_03	KS22J6
			14392	KX513732	<i>T. sp.</i> ‘black’	Ht_04	KS23A10
			14452	KX513673	<i>T. sp.</i> ‘black’	Ht_25	T14/03/A2
			14453	KX513672	<i>T. sp.</i> ‘black’	Ht_26	T14/03/A3
				Z12096	<i>T. sp.</i> ‘black’	Ht_03	
				Z12094	<i>T. sp.</i> ‘black’	Ht_03	
Bulumbu	−3,77	29,12	14387	KX513737	<i>T. sp.</i> ‘black’	Ht_01*	KS22J6
			816	KX513597	<i>T. sp.</i> ‘black’	Ht_02*	816
			820	KX513596	<i>T. sp.</i> ‘black’	Ht_02*	820
			833	KX513604	<i>T. sp.</i> ‘black’	Ht_51	833
Caramba	−4,5	29,18	835	KX513595	<i>T. sp.</i> ‘black’	Ht_15*	835
			14965	KX513738	<i>T. sp.</i> ‘black’	Ht_59	T14/05/A6
Kabezi	−3,5	29,333		Z75702	<i>T. sp.</i> ‘black’	Ht_48	
			14966	KX513739	<i>T. sp.</i> ‘black’	Ht_60*	T14/05/A7
				Z75694	<i>T. sp.</i> ‘black’	Ht_47	
Kagongo	−3,7352	29,5847	14415	KX513709	<i>T. sp.</i> ‘black’	Ht_15	KS20I9
			14502	KX513624	<i>T. sp.</i> ‘black’	Ht_15	T14/03/F2
			14416	KX513708	<i>T. sp.</i> ‘black’	Ht_16	KS20I9
			14427	KX513698	<i>T. sp.</i> ‘black’	Ht_16	KS20I5
			14432	KX513693	<i>T. sp.</i> ‘black’	Ht_16	KS20I6
			14495	KX513631	<i>T. sp.</i> ‘black’	Ht_16	T14/03/E5

Table 1 continued

Location	Latitude (S)	Longitude (E)	Extraction no.	Accession no.	Species	Haplotype	Sample ID
Kalundu	−3,388	29,144	14498	KX513628	<i>T. sp.</i> ‘black’	Ht_16	T14/03/E8
			14418	KX513706	<i>T. sp.</i> ‘black’	Ht_17	KS20I9
			14496	KX513630	<i>T. sp.</i> ‘black’	Ht_17	T14/03/E6
			14501	KX513625	<i>T. sp.</i> ‘black’	Ht_17	T14/03/F1
			14503	KX513623	<i>T. sp.</i> ‘black’	Ht_17	T14/03/F3
			14431	KX513694	<i>T. sp.</i> ‘black’	Ht_18	KS20I5
			14434	KX513691	<i>T. sp.</i> ‘black’	Ht_19	KS20I6
			14497	KX513629	<i>T. sp.</i> ‘black’	Ht_40	T14/03/E7
			14417	KX513707	<i>T. sp.</i> ‘black’	Ht_16	KS20I9
			14428	KX513697	<i>T. sp.</i> ‘black’	Ht_16	KS20I5
			14429	KX513696	<i>T. sp.</i> ‘black’	Ht_16	KS20I5
			14430	KX513695	<i>T. sp.</i> ‘black’	Ht_16	KS20I5
			14433	KX513692	<i>T. sp.</i> ‘black’	Ht_16	KS20I6
			14494	KX513632	<i>T. sp.</i> ‘black’	Ht_15	T14/03/E4
			14499	KX513627	<i>T. sp.</i> ‘black’	Ht_16	T14/03/E9
			14500	KX513626	<i>T. sp.</i> ‘black’	Ht_16	T14/03/E10
			14493	KX513633	<i>T. sp.</i> ‘black’	Ht_15*	T14/03/E3
			14492	KX513634	<i>T. sp.</i> ‘black’	Ht_16*	T14/03/E2
			14438	KX513687	<i>T. sp.</i> ‘black’	Ht_01	KS24G3
			14476	KX513650	<i>T. sp.</i> ‘black’	Ht_10	T14/03/C6
			14436	KX513689	<i>T. sp.</i> ‘black’	Ht_11	KS24G3
			14439	KX513686	<i>T. sp.</i> ‘black’	Ht_11	KS24G3
			14473	KX513653	<i>T. sp.</i> ‘black’	Ht_11	T14/03/C3
			14478	KX513648	<i>T. sp.</i> ‘black’	Ht_11	T14/03/C8
			14481	KX513645	<i>T. sp.</i> ‘black’	Ht_11	T14/03/D1
			14441	KX513684	<i>T. sp.</i> ‘black’	Ht_14	KS24G4
			14474	KX513652	<i>T. sp.</i> ‘black’	Ht_14	T14/03/C4
			14437	KX513688	<i>T. sp.</i> ‘black’	Ht_20	KS24G3
			14442	KX513683	<i>T. sp.</i> ‘black’	Ht_21	KS24G4
			14475	KX513651	<i>T. sp.</i> ‘black’	Ht_21	T14/03/C5
			14472	KX513654	<i>T. sp.</i> ‘black’	Ht_30	T14/03/C2
			14479	KX513647	<i>T. sp.</i> ‘black’	Ht_35	T14/03/C9
			14477	KX513649	<i>T. sp.</i> ‘black’	Ht_38	T14/03/C7
			14480	KX513646	<i>T. sp.</i> ‘black’	Ht_35	T14/03/C10
			14435	KX513690	<i>T. sp.</i> ‘black’	Ht_01*	KS24G3
			14440	KX513685	<i>T. sp.</i> ‘black’	Ht_11*	KS24G4
Kiriza	−4,05	29,216		Z75700	<i>T. sp.</i> ‘black’	Ht_16	
				Z12070	<i>T. sp.</i> ‘black’	Ht_55	
Kisokwe	−4,24	29,18	890	KX513593	<i>T. sp.</i> ‘black’	Ht_53*	890
Lubumba	−3,52	29,15	865	KX513594	<i>T. sp.</i> ‘black’	Ht_52	865
Luhanga	−3,5186	29,3969	14403	KX513721	<i>T. sp.</i> ‘black’	Ht_01	KS24B8
			14456	KX513669	<i>T. sp.</i> ‘black’	Ht_01	T14/03/A6

Table 1 continued

Location	Latitude (S)	Longitude (E)	Extraction no.	Accession no.	Species	Haplotype	Sample ID
Magara	−3,726	29,31	14405	KX513719	<i>T. sp.</i> ‘black’	Ht_09	KS24B8
			14406	KX513718	<i>T. sp.</i> ‘black’	Ht_10	KS24B8
			14407	KX513717	<i>T. sp.</i> ‘black’	Ht_11	KS24B8
			14457	KX513668	<i>T. sp.</i> ‘black’	Ht_11	T14/03/A7
			14458	KX513667	<i>T. sp.</i> ‘black’	Ht_27	T14/03/A8
			14459	KX513666	<i>T. sp.</i> ‘black’	Ht_28	T14/03/A9
			14460	KX513665	<i>T. sp.</i> ‘black’	Ht_29	T14/03/A10
			14462	KX513663	<i>T. sp.</i> ‘black’	Ht_29	T14/03/B2
			14461	KX513664	<i>T. sp.</i> ‘black’	Ht_30	T14/03/B1
			14467	KX513658	<i>T. sp.</i> ‘black’	Ht_30	T14/03/B7
			14463	KX513662	<i>T. sp.</i> ‘black’	Ht_31	T14/03/B3
			14464	KX513661	<i>T. sp.</i> ‘black’	Ht_32	T14/03/B4
			14465	KX513660	<i>T. sp.</i> ‘black’	Ht_33	T14/03/B5
			14466	KX513659	<i>T. sp.</i> ‘black’	Ht_34	T14/03/B6
			14468	KX513657	<i>T. sp.</i> ‘black’	Ht_35	T14/03/B8
			14470	KX513605	<i>T. sp.</i> ‘black’	Ht_35	T14/03/B10
			14469	KX513656	<i>T. sp.</i> ‘black’	Ht_36	T14/03/B9
			14471	KX513655	<i>T. sp.</i> ‘black’	Ht_37	T14/03/C1
			14404	KX513720	<i>T. sp.</i> ‘black’	Ht_01	KS24B8
			14408	KX513716	<i>T. sp.</i> ‘black’	Ht_11	KS24D6
			14409	KX513715	<i>T. sp.</i> ‘black’	Ht_11	KS24D6
			14410	KX513714	<i>T. sp.</i> ‘black’	Ht_11	KS24D6
			783	KX513602	<i>T. sp.</i> ‘black’	Ht_27	783
			784	KX513601	<i>T. sp.</i> ‘black’	Ht_27	784
			787	KX513598	<i>T. sp.</i> ‘black’	Ht_09	787
			785	KX513600	<i>T. sp.</i> ‘black’	Ht_11*	785
			786	KX513599	<i>T. sp.</i> ‘black’	Ht_27*	786
			14395	KX513729	<i>T. sp.</i> ‘black’	Ht_05	KS20G7
			14486	KX513640	<i>T. sp.</i> ‘black’	Ht_05	T14/03/D6
			14396	KX513728	<i>T. sp.</i> ‘black’	Ht_06	KS20G7
			14401	KX513723	<i>T. sp.</i> ‘black’	Ht_06	KS20G8
			14483	KX513643	<i>T. sp.</i> ‘black’	Ht_06	T14/03/D3
			14485	KX513641	<i>T. sp.</i> ‘black’	Ht_06	T14/03/D5
			14490	KX513636	<i>T. sp.</i> ‘black’	Ht_06	T14/03/D10
			14397	KX513727	<i>T. sp.</i> ‘black’	Ht_07	KS20G7
			14399	KX513725	<i>T. sp.</i> ‘black’	Ht_07	KS20G7
			14484	KX513642	<i>T. sp.</i> ‘black’	Ht_07	T14/03/D4
			14488	KX513638	<i>T. sp.</i> ‘black’	Ht_07	T14/03/D8
			14491	KX513635	<i>T. sp.</i> ‘black’	Ht_07	T14/03/E1
			14398	KX513726	<i>T. sp.</i> ‘black’	Ht_08	KS20G7
			14402	KX513722	<i>T. sp.</i> ‘black’	Ht_08	KS20G8
			14487	KX513639	<i>T. sp.</i> ‘black’	Ht_39	T14/03/D7
			14489	KX513637	<i>T. sp.</i> ‘black’	Ht_39	T14/03/D9
			14400	KX513724	<i>T. sp.</i> ‘black’	Ht_07	KS20G8

Table 1 continued

Location	Latitude (S)	Longitude (E)	Extraction no.	Accession no.	Species	Haplotype	Sample ID
Mboko	−3,9166	29,083	14482	KX513644	<i>T. sp. 'black'</i>	Ht_07*	T14/03/D2
				AY660763	<i>T. sp. 'black'</i>	Ht_50	
				Z75700	<i>T. sp. 'black'</i>	Ht_16	
			14509	KX513618	<i>T. brichardi</i>	Ht_12	T14/03/F8
			14517	KX513611	<i>T. brichardi</i>	Ht_12	T14/03/G6
			14510	KX513617	<i>T. brichardi</i>	Ht_12	T14/03/F9
			14511	KX513616	<i>T. brichardi</i>	Ht_12	T14/03/F10
			14512	KX513615	<i>T. brichardi</i>	Ht_12	T14/03/G1
			14513	KX513614	<i>T. brichardi</i>	Ht_12	T14/03/G2
			14514	KX513613	<i>T. brichardi</i>	Ht_12	T14/03/G3
			14443	KX513682	<i>T. brichardi</i>	Ht_12	KS25E2
			14445	KX513680	<i>T. brichardi</i>	Ht_12	KS25E2
			14449	KX513676	<i>T. brichardi</i>	Ht_12	KS25G6
			14450	KX513675	<i>T. brichardi</i>	Ht_13	KS25G6
			14505	KX513621	<i>T. brichardi</i>	Ht_22	T14/03/F5
			14520	KX513608	<i>T. brichardi</i>	Ht_22	T14/03/G9
			14444	KX513681	<i>T. brichardi</i>	Ht_22	KS25E2
			14447	KX513678	<i>T. brichardi</i>	Ht_23	KS25E3
			14518	KX513610	<i>T. brichardi</i>	Ht_24	T14/03/G7
			14448	KX513677	<i>T. brichardi</i>	Ht_24	KS25G6
Mvugo	−3,9808	29,503	14504	KX513622	<i>T. brichardi</i>	Ht_41	T14/03/F4
			14519	KX513609	<i>T. brichardi</i>	Ht_41	T14/03/G8
			14507	KX513619	<i>T. brichardi</i>	Ht_42	T14/03/F7
			14515	KX513612	<i>T. brichardi</i>	Ht_43	T14/03/G4
			14521	KX513607	<i>T. brichardi</i>	Ht_44	T14/03/G10
			14506	KX513620	<i>T. brichardi</i>	Ht_22	T14/03/F6
			14446	KX513679	<i>T. brichardi</i>	Ht_12	KS25E2
			14411	KX513713	<i>T. brichardi</i>	Ht_12	KS25H7
			14412	KX513712	<i>T. brichardi</i>	Ht_13	KS25H7
Ngombe	−4,666	29,6166		AJ489622	<i>T. brichardi</i>	Ht_01	
				AJ295923	<i>T. brichardi</i>	Ht_57	
				AJ95924	<i>T. brichardi</i>	Ht_58	
Nyahurongoka	−3,69	29,33	14414	KX513710	<i>T. sp. 'black'</i>	Ht_06	KS20A2
			14413	KX513711	<i>T. sp. 'black'</i>	Ht_14	KS20I2
			1386	KX513603	<i>T. sp. 'black'</i>	Ht_07	1386
			1387	KX513592	<i>T. sp. 'black'</i>	Ht_54	1387
Nyanza Lac	−4,33	29,583	14969	KX513742	<i>T. brichardi</i>	Ht_46	KS32B1
				Z12054	<i>T. brichardi</i>	Ht_46	
			14968	KX513741	<i>T. brichardi</i>	Ht_46*	KS32A9
Rutunga	−3,666	29,316	14967	KX513740	<i>T. brichardi</i>	Ht_46*	KS32A8
			14522	KX513606	<i>T. sp. 'black'</i>	Ht_45	T14/03/H1

Table 1 continued

Location	Latitude (S)	Longitude (E)	Extraction no.	Accession no.	Species	Haplotype	Sample ID
'Ubwari-green'	−4,164	29,2582	13859	Z12050	<i>T. sp.</i> 'black'	Ht_49	T13/1/B1
				Z12051	<i>T. sp.</i> 'black'	Ht_06	
				Z12049	<i>T. sp.</i> 'black'	Ht_49	
				AY660840	<i>T. cf. brichardi</i>	Ht_56	
				AY660842	<i>T. sp.</i> 'black'	Ht_56	
				AY660843	<i>T. sp.</i> 'black'	Ht_56	
Ubwari Peninsula (color morph not known)				KX513591	<i>T. sp.</i> 'black'	Ht_56*	T13/1/B1
				AY660841	<i>T. sp.</i>	Ht_48	

Individuals marked in bold letters and asterisks were also used in the AFLP tree

of a larger dataset (Van Steenberg, unpublished data) in which negative controls and a minimum of 19 replicates per primer combination were included in the reactions. Size and peak height of fragments between 100 and 500 bp were determined using GeneMapper v.3.7 (Applied Biosystems). Bins were checked by eye and preprocessed for threshold optimization for locus retention and phenotype calling with AFLP-SCORE 1.4a (Whitlock et al., 2008) following (Mattersdorfer et al., 2012). The average mismatch error rate was 1.26%. The final binary matrix for the representative set of northern *Tropheus* consisted of 442 polymorphic characters.

Phylogenetic analysis

Phylogenetic relationships among mitochondrial haplotypes were inferred by means of a neighborjoining (NJ) tree in MEGA 6.0 (Tamura et al., 2013). For NJ tree inference, identical sequences were collapsed into haplotypes using DnaSP 5.10 (Rozas, 2010). Based on the Bayesian information criterion (BIC), jModelTest 0.1 (Posada, 2008) identified the HKY+G (Hasegawa et al., 1985a, b) model as the best fitting model of molecular evolution. As this model is not implemented in MEGA, the TN93+G (Tamura & Nei, 1993) model—as the best fitting model available in MEGA—was employed for NJ tree inference instead. Nodal support was assessed by means of bootstrapping (10,000 pseudoreplicates). Furthermore, a statistical parsimony network (Templeton et al., 1992) was constructed in POPART (Leigh and Bryant, 2015).

To get an idea about the putative timing of major divergence events among and within the main

mitochondrial lineages, we inferred a time-calibrated mitochondrial tree in BEAST 1.8 (Drummond & Rambaut, 2007). Two independent MCMC chains were run for 10^6 generations, with model parameters and trees sampled every 1000 generations. We employed the HKY+G substitution model with a strict molecular clock (as we are looking at mostly intraspecific data; Brown & Yang, 2011) assuming minimum and maximum substitution rates of 3.24 and 5.7% per MY, respectively (Koblmüller et al., 2009; Genner et al., 2010), and a Bayesian skyline tree prior (Drummond et al., 2005). All other priors were left at default. The first 10% of generations were discarded from each log and tree file as burn-in before the two chains were combined using LogCombiner (available as part of the BEAST package). Chain convergence to stationarity for all model parameters was assessed in Tracer 1.6 (available from <http://beast.bio.ed.ac.uk/tracer>). The pooled post-burn-in Effective Sample Sizes (ESS) for all parameters exceeded 200, indicating that the pooled log file accurately represented the posterior distribution (Kuhner, 2009). Divergence times were derived from the pooled post-burn-in results and TreeAnnotator (available as part of the BEAST package) was used to compute a maximum-clade-credibility tree, which was visualized in FigTree 1.4.1 (available from <http://beast.bio.ed.ac.uk/figtree>). Divergence times were calculated as median node heights of the 95% highest posterior density (HPD) intervals.

HEXt (Schneider et al., 2016) was used to infer a NJ tree from the AFLP data based on Nei-Li distances (Nei & Li, 1979) and estimate statistical support from 1000 bootstrap replicates. The tree was rooted with

Tropheus duboisi, which was previously shown to represent the sister group of all other *Tropheus* (Koblmüller et al., 2010).

Within- and among-population patterns of genetic diversity

Genetic diversity indices—number of haplotypes (H), Haplotype diversity (H_d), nucleotide diversity (π)—as well as mismatch distributions were calculated in DnaSP for all populations with a sample size $N \geq 14$. Population differentiation was estimated by θ_{ST} (Weir & Cockerham, 1984) and Φ_{ST} (Excoffier et al., 1992) in Arlequin v.3.5 (Excoffier & Lischer, 2010), with significance inference corrected for multiple testing following (Benjamini & Hochberg, 1995).

Past population size trajectories for main mitochondrial lineages and populations (with sample size $N \geq 9$) were inferred by means of Bayesian skyline

plots (BSPs; Drummond et al., 2005) in BEAST and visualized in Tracer, employing the same settings as for the time-calibrated mitochondrial tree (see above). The various datasets required different run lengths, but all analyses were run until ESS for all parameters were >200 .

Results

Phylogeographic patterns

The combined analysis of 179 individuals from 21 localities in Congo and Burundi identified 60 haplotypes and corroborates the assignment of all *Tropheus* from this area to two major mtDNA lineages (Fig. 2), TCS-lineages 1-A and 2-B (following Sturmhuber et al., 2005). Moreover, each of the mitochondrial clusters comprises two to three subclusters, coded in

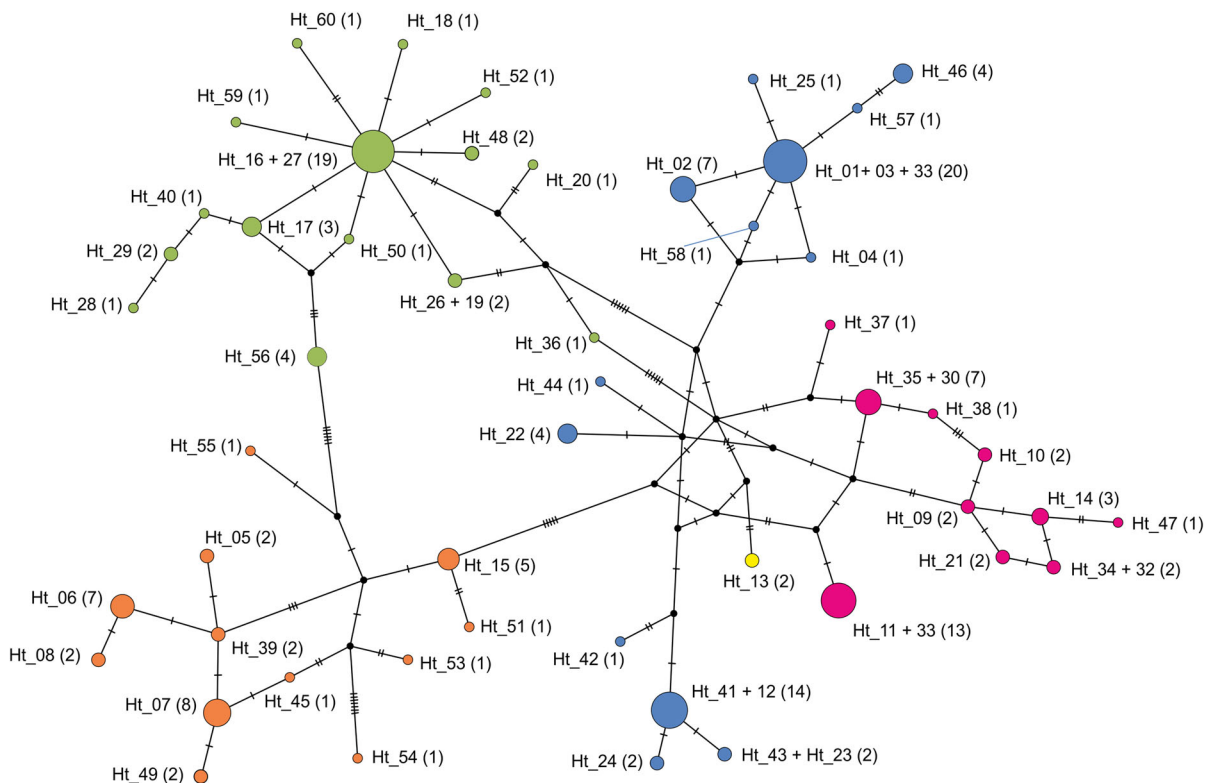


Fig. 2 Statistical parsimony network of northern *Tropheus* haplotypes based on a 352 bp long segment of the mitochondrial control region. Each haplotype is represented by a circle, the size of which correlates with the number of individuals sharing the same haplotype. Small bars indicate the number of

substitutions between haplotypes. Each of the mitochondrial clusters comprises two to three subclusters, coded in yellow, blue and pink for TCS-lineage 1, and in green and red for TCS-lineage 2 in all figures

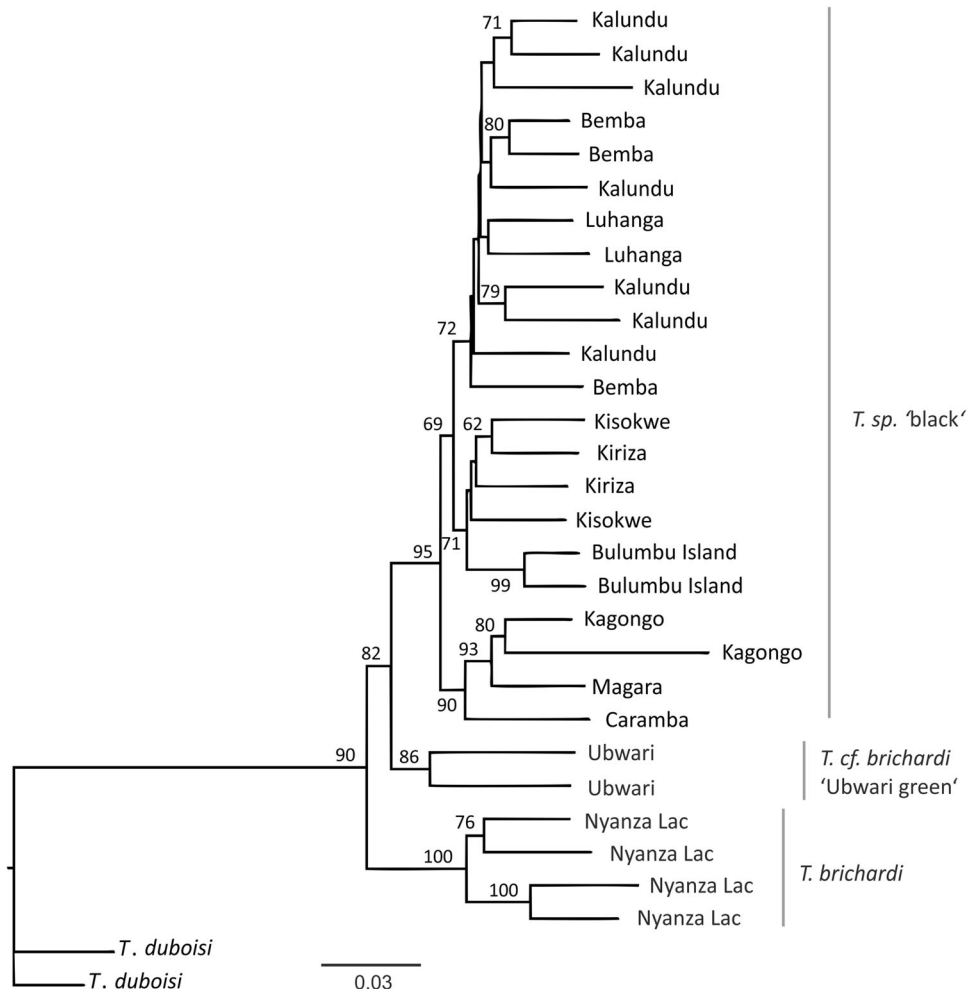


Fig. 3 NJ tree based on 442 polymorphic AFLP loci. Only bootstrap values >50 are shown. *Tropheus duboisi* was used as outgroup

yellow, blue and pink for TCS-lineage 1, and in green and red for TCS-lineage 2 in all Figures. While the mtDNA tree is partially inconsistent with the current species assignment, the AFLP-based nuclear tree shows the reciprocal monophyly of the two *Tropheus* species present in the northernmost part of the Lake, *T. brichardi* and *T. sp. 'black'*, in relation to the outgroup *T. duboisi* (Fig. 3). While the individuals of *T. sp. 'black'* Caramba are resolved in the clade of all other *T. sp. 'black'*, the phenotypically aberrant *Tropheus* cf. *brichardi* 'Ubwari-green' from Cape Muzimu southwards occupy an intermediate position in the AFLP-tree (Fig. 3, note that one previously published sample from the Ubwari Peninsula was not explicitly assigned to either of the two taxonomic entities; it is thus labeled as *T. sp. Ubwari Peninsula* in Table 1

and the figures). The time-calibrated mitochondrial tree (Fig. 4) suggests that TCS-lineages 1 and 2 diverged roughly 450–800 KYA and indicates simultaneous east/west divergence in three out of the five mitochondrial sublineages about 150–265 KYA. The presence of shared or closely related haplotypes on opposite shores (in the yellow sublineage the sample size was too small) indicated gene flow between eastern and western populations in the more recent past. Several populations comprised haplotypes pertaining to different haplogroups (see haplogroup distributions along shoreline in Fig. 5, and mismatch distributions in Supplementary Fig. 1). Concerning the distribution of haplotype lineages, one can clearly observe that along the eastern shoreline northwards the distribution of populations with blue and pink

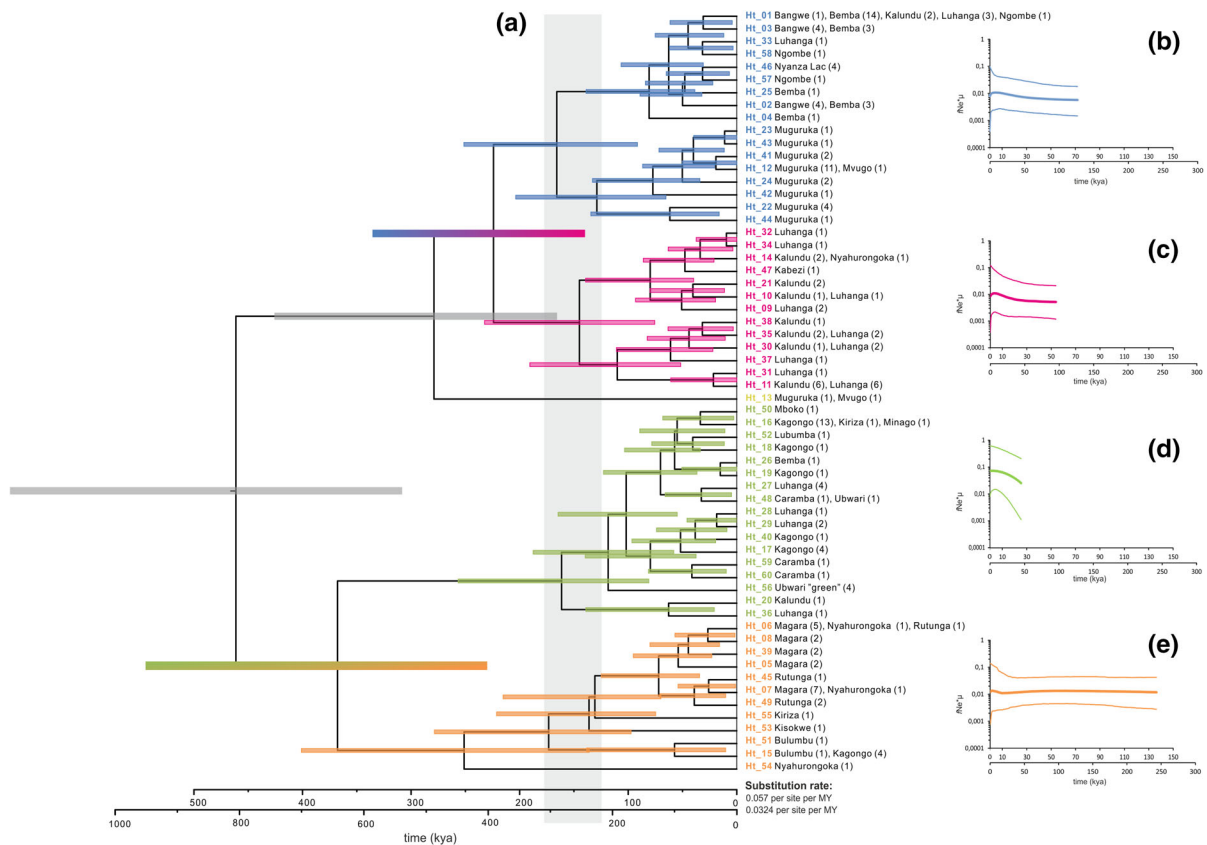


Fig. 4 Chronogram of the diversification of northern *Tropheus* based on mitochondrial control region haplotypes. Divergence time estimates are represented as the median node height of the 95% highest posterior density (HPD) interval from a BEAST maximum-clade-credibility tree. Node bars span the 95% HPD interval for each node. The colors indicate major haplotype lineages comprising two to three subclusters, coded in yellow,

haplotypes (TCS-1) is interrupted by populations with green and orange haplotypes (TCS-2), and that blue and pink haplotypes (TCS-1) again dominate the very north of the lake. The green and orange TCS-2 haplotypes, however, dominate the western shores at the Ubwari Peninsula, to be progressively replaced northwards by blue TCS-1 haplotypes at Bemba/Bangwe and blue/pink haplotypes at Luhanga/Kalundu. From the distribution of identical and very closely related haplotypes (1–2 mutations), one can deduce recent long-distance dispersal of blue haplotypes from Nyanza Lac to the Bemba-Kalundu stretch, as well as intense connections of green and orange haplotypes across the ridge at the eastern and western Ubwari Faults (Fig. 5). The two sympatric *Tropheus*, *T. sp.* ‘black’ Caramba and *T. cf. brichardi* ‘Ubwari-

blue, and pink for TCS-lineage 1, and in green and red for TCS-lineage 2 (Fig. 2). Insets to the right b–e show past population size changes for the four major mitochondrial clades, as inferred by means of Bayesian skyline plots. The y-axis represents the population size parameter (female effective population size times the mutation rate). Thick and thin lines denote median estimates and 95% HPD intervals, respectively

green’, share the same mtDNA lineage (TCS-2), despite their different positions in the AFLP tree (Fig. 3), suggesting hybridization and subsequent mitochondrial capture. The BSPs revealed signatures of Late Pleistocene population growth in all but the orange subcluster (Fig. 4). Particularly strong population expansion is evident for the green subcluster. Whether the recent population size decline (over last few hundreds of years) apparent in most mitochondrial lineages is a true signal potentially correlated with increasing human population sizes along the lake shore or simply a methodological artifact (Chikhi et al., 2010; Heller et al., 2013) remains unclear.

It is also interesting to note that the TCS-1 haplotypes assigned to the pink haplotype cluster are not exclusive to the very northern part of the lake. This

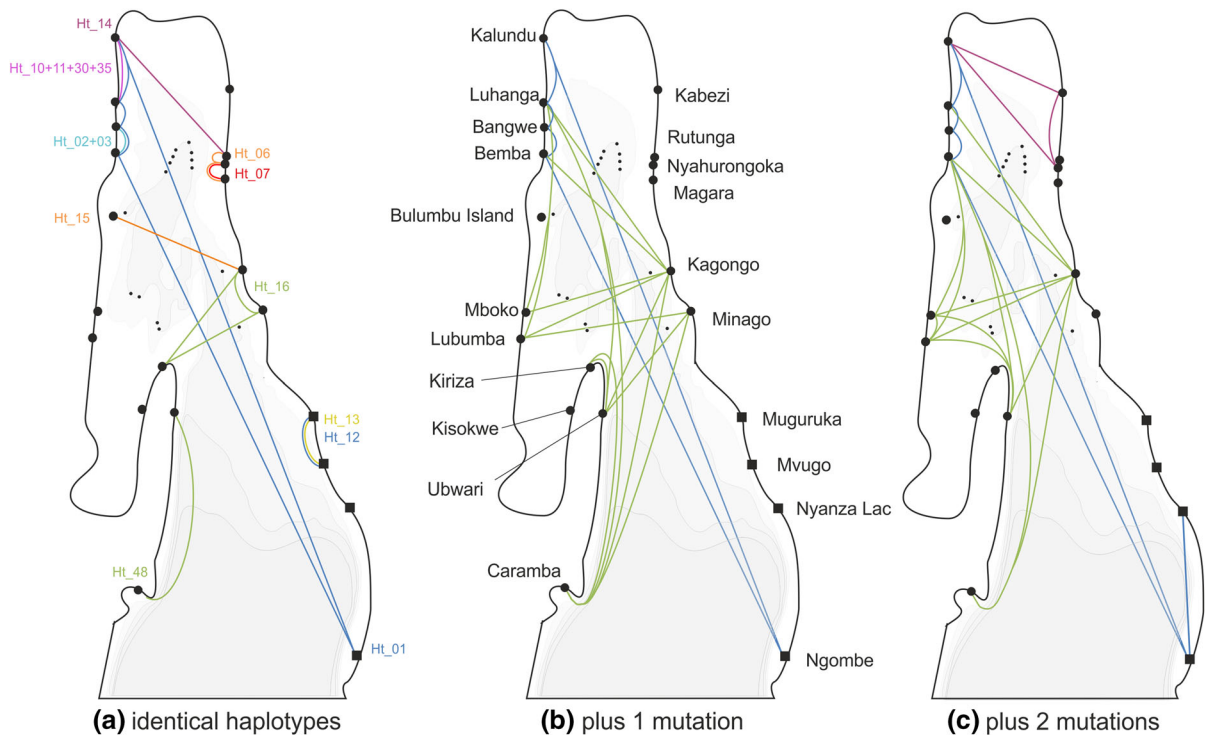


Fig. 5 Distribution of identical and similar (1–2 mutations difference) *Tropheus* haplotypes in northern Lake Tanganyika. Colors are assigned to the clades as described in the text.

haplogroup is also found in some individuals of *T. moorii* from the southern basin such at Fulwe and Wapembe, as well as in fish at Kasakalawe from the very southern end of the lake. In-between, we only have a record of the pink haplotype cluster in a single *T. sp.* ‘black’ Kirschfleck individual from Mabilibili at the central eastern shore. These biogeographic data suggest a particularly widespread migration of the members of this TCS-sublineage (Sturmbauer et al., 2005).

Population genetics

The number of haplotypes per population varied considerably among populations, with the lowest number detected in Magara ($N = 5$) and the highest number detected in Luhanga ($N = 11$). Estimates of haplotype diversity (H_d) and nucleotide diversity (π) ranged from 0.674 (Kagongo) to 0.897 (Luhanga) and 0.00406 (Magara) to 0.02420 (Luhanga) (Table 2). The large variation of π across populations indicates that different numbers of haplogroups contributed to the genetic diversity in the different populations.

Squares marking sampling sites indicate the occurrence of *T. brichardi* instead of *T. sp.* ‘black’ (circle symbols)

Population genetic differentiation was high and highly significant for most pairwise comparisons. Only the two northernmost populations at the western shoreline were not significantly differentiated from each other (Table 2). The BSPs revealed clear signatures of simultaneous recent population growth for four populations (Bemba, Bangwe, Magara, Kagongo), whereas the three other populations (Luhanga, Kalundu, Muguruka) appear to have experienced a decline in the recent past (Fig. 6).

Discussion

Phylogeography

Our new data fill important gaps and allow us to hypothesize a colonization and admixis scenario for *Tropheus* populations inhabiting the very northern section of Lake Tanganyika, and highlight the combined effect of recurrent drastic lake level fluctuations and a complex basin structure on the phylogeographic structure of rock-dwelling cichlids in this part of the

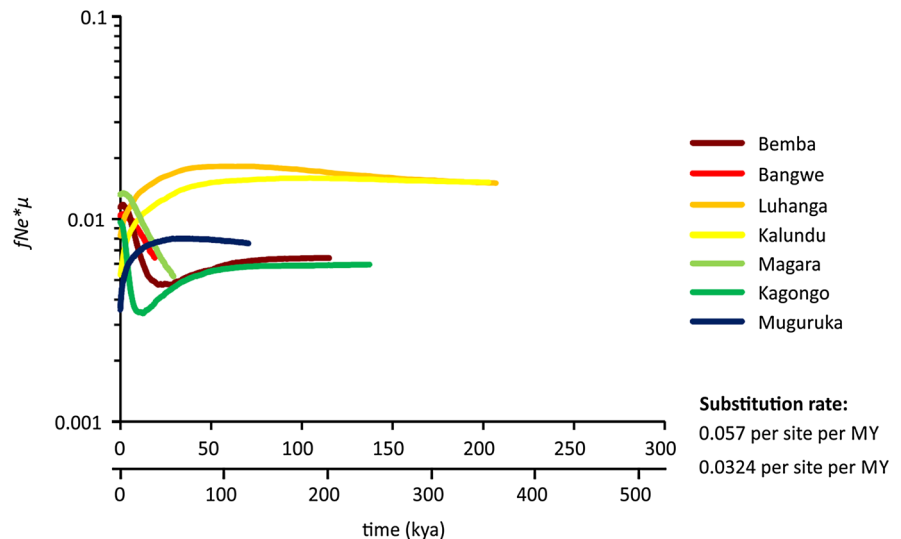
Table 2 Population sample sizes (N), genetic diversity and pairwise population genetic differentiation based on the most variable part of the mitochondrial control region

	N	H	H_d	π	Bemba	Luhanga	Kalundu	Magara	Kagongo	Muguruka
Bemba	14	6	0.736	0.00652		0.113***	0.144**	0.242***	0.298***	0.244***
Luhanga	29	11	0.897	0.02420	0.311***		0.002	0.142***	0.194***	0.147***
Kalundu	18	8	0.863	0.01634	0.489***	0.037		0.173***	0.230***	0.178***
Magara	18	5	0.778	0.00406	0.882***	0.625***	0.755***		0.277***	0.225***
Kagongo	24	6	0.674	0.01114	0.683***	0.368***	0.605***	0.785***		0.277***
Muguruka	24	7	0.688	0.01151	0.676***	0.419***	0.504***	0.844***	0.757***	

Only populations with $N \geq 14$ are included

H number of haplotypes, H_d gene diversity, π nucleotide diversity. Above diagonal: θ_{ST} estimates, below diagonal: Φ_{ST} values. Significance levels, $P < 0.05$, < 0.01 and < 0.001 , after correction for multiple test, are indicated as *, **, and ***, respectively

Fig. 6 Bayesian skyline plots of population sizes through time for populations with a sample size $N \geq 9$. Depicted are the median estimates. The y-axis represents the population size parameter (female effective population size times the mutation rate)



lake (Cohen et al., 2007). This region comprises a small subbasin at minus 300 m north of the Ubwari Penninsula, named the Bujumbura basin, separated by a ridge from the Kigoma basin. This basin played an important role as lowstand refugium and melting pot for two mtDNA lineages of *Tropheus* in the more recent past. Our data suggest that it seems most likely that the original *Tropheus* population of the Bujumbura basin comprised *T. sp.* 'black,' given that all populations inhabiting this basin are assigned to this species. The original *T. sp.* 'black' population was substantially perturbed by a series of lake level fluctuations, triggering the invasion of *T. brichardi* from the Kigoma basin along the eastern shoreline. At the eastern shore of the Bujumbura basin the present-day distribution of identical or closely related mtDNA

haplotypes assigned to the *T. brichardi* lineage (TCS-1) is discontinuous. As the pink TCS-1 haplotype sublineage is exclusively distributed in the very north of the Bujumbura basin and shows great diversity only there, while the blue TCS-1 haplotype sublineage is equally diverse but much more widespread, ranging from the very northwest of the present-day lake down to Muguruka and even Ngombe, we suggest that it is more parsimonious to assume (at least) two migration waves for the *T. brichardi*-lineage. The members of the pink TCS-1 haplotypes are likely to have arrived at the first wave. In-between the two migration waves, during another period of lower lake level, *T. sp.* 'black' with TCS-2 haplotypes crossed over from the west along the Ubwari ridge and largely replaced the *T. brichardi*-like TCS-1 populations from Minago to

Magara. That their colonization of the eastern shores between Minago and Magara is not very recent can be delineated from the evolution of two distinct TCS-2 haplotype subgroups, one with its center of diversity at the eastern and the other at the western shore. At least one subsequent re-connection blurred the distribution pattern of TCS-2 haplotypes and allowed large-scale admixis with TCS-1 haplotypes from the blue sublineage. That this admixis event was very recently, probably during the last glacial maximum, can be seen from the long-distance distribution of identical and very closely related haplotypes in both lineages (Fig. 5). This (re)colonization by *T. brichardi* might have been successful due to a local deterioration of the habitat. Both the volcanic activity in the Bemba area (Pflumio et al., 1994) and the erratic flow of the sediment carrying Rusizi River (Cassanova & Hillaire-Marcel, 1992) could be responsible for this.

Albeit all populations of the Bujumbura basin are *T. sp.* ‘black,’ despite of the complex phylogeographic pattern of TCS-1 and TCS-2 haplotypes, many seem heavily introgressed by *T. brichardi* TCS-1 haplotypes and sometimes phenotypically intermediate (Van Steenberge, 2014). This is also supported by the presence of TCS-1 haplotypes at Nyahurongoka and Kabezi. Since the rock shores around the Ubwari Peninsula are dominated by TCS-2 haplotypes, despite the partial sympatry of two species, we suggest that this section represents their original distribution area, and thus the original distribution area of *T. sp.* ‘black.’ It seems likely that the TCS-2 haplotypes sampled from Minago to Rutunga, as they interrupt the distribution of TCS-1 haplotypes there, made it across the bottleneck area northeast of Cape Banza between the West and East Ubwari Faults. The lowstand-shoreline in this area over the ridge is extremely narrow and resembles a meandric river, so that complex admixis scenarios along both basin edges are inevitable. This may have allowed repeated bridging of mainly TCS-2 haplotypes (see Fig. 1 in Lezzar et al., 2002). A complex colonization-admixis-partial extinction scenario is also supported by the present-day distribution of the more distantly related *Tropheus duboisi*, which occurs almost continuously together with TCS-1 haplotypes of *T. brichardi*-provenience at the east coast from the Mahale Mountains to Muguruka, but only occurs at a single small stretch at the NW coast near Bemba, together with a TCS-1 dominated *Tropheus sp.* ‘black’

population sharing identical haplotypes with their allies at Ngombe. In fact, the control region haplotypes in the *T. duboisi* populations from Bemba and Mwamugongo (near Ngombe) are separated by 2 mutations only (Van Steenberge et al., 2015). Moreover, *Julidochromis* species show similar cross-bottleneck distribution, in that *J. regani* shares the habitat with “pure” *T. brichardi* around Nyanza Lac, followed northwards by a putative hybrid between *J. regani* and *J. marlieri* (*J. regani affinis* from Rumonge northwards, as well as at the tip of the Ubwari Peninsula), followed by pure *J. marlieri* around Rutunga, and finally followed by pure *J. regani* at Bujumbura again (Brichard, 1978).

Concerning the lake-wide distribution patterns of the TCS-1 haplotype lineage and its subclusters, it is important to note that the pink TCS-1 subcluster is very widespread but rare (Sturmbauer et al., 2005). Aside of the very North of the lake it only occurs in a single individual at Mabilibili, in populations at Fulwe and north of Wapembe, and at Kasakalawe at the very southern end of the lake. This distribution points to a large migration wave that was overlaid by subsequent migration waves and masked by lineage sorting in most areas. The survival of this haplotype subgroup at both ends of the lake might point to a lesser chance to be overlaid at such tip-populations. The observed widespread but rare distribution pattern of the pink TCS-1 subcluster in fact supports our suggestion of at least two arrivals of TCS-1 haplotype fish in the very North of the Lake.

Taxonomic considerations

It is important to note that the distribution of mtDNA haplotypes, in combination with nuclear DNA markers, morphology and color, indicates considerable levels of past gene flow among entities considered as valid biological species. It was suggested that massive perturbations of the habitat or availability of new habitat cause species to interbreed even if they would not do so under stable conditions (Rüber et al., 2001; Seehausen, 2004). This scenario seems to fit many of the here studied *Tropheus* populations, as we found evidence for extensive mitochondrial introgression, which did not become evident in the nuclear data obtained so far.

Under particular circumstances hybridization upon secondary contact might actually lead to a novel

evolutionary stable entity, as, for example, suggested for a number of animal taxa (Grant & Grant, 2002; Gompert et al., 2006; Larsen et al., 2010; Seft & Koblmüller, 2016), including other cichlid fishes (Salzburger et al., 2002). It has been shown that hybrid populations can rapidly generate novel (“transgressive”) phenotypes which might be rapidly sorted out via natural selection, or alternatively, that hybrid populations become isolated from their sources in particular habitats, to form a novel entity distinct from both parental species (Salzburger et al., 2002; Parsons et al., 2011). Also in *Tropheus*, the importance of hybridization for generating novel phenotypes has been proposed previously (Egger et al., 2007). Among the *Tropheus* morphs included here, *Tropheus* cf. *brichardi* ‘Ubwari-green’, exhibits an intermediate phenotype in form of greenish *T. brichardi*-like body color, blue eyes, but red-yellow *T. sp.* ‘black’ banding on the body flanks. In the mitochondrial data, all *Tropheus* cf. *brichardi* Ubwari-green samples analyzed group among *T. sp.* ‘black,’ whereas the AFLP-based nuclear DNA assignment also suggests intermediacy (Fig. 3). It is important to repeat here that this *Tropheus* lives in full sympatry with *T. sp.* ‘black’ Caramba at one particular shoreline at the western side of the Ubwari Peninsula between Cape Caramba and Muzimu (Brichard, 1978; Konings, 2013), so that speciation can be assumed as completed in these populations. The two species are not segregating spatially like many other sympatric *Tropheus* but fully coexist in similar abundancies (Ad Konings personal communication). As *Tropheus* cf. *brichardi* ‘Ubwari-green’ combines features of both parental species and lives in sympatry with one of them, it clearly cannot be assigned to either of the described species, and should be regarded as a distinct species, for which hybrid origin seems likely.

Concerning taxonomic assignments, the populations from Ngombe to Muguruka should represent “pure” *Tropheus brichardi*, as these exclusively comprise TCS-1 haplotypes and because of phenotypic similarities to specimens collected in Nyanza Lac, the type locality. Unfortunately, no AFLP data are available for localities other than Nyanza Lac to support our species assignment also by nuclear multilocus data. All other populations except for *T. cf. brichardi* ‘Ubwari-green’ are pure *T. sp.* ‘black.’ Even if some populations show evidence for past mitochondrial introgression from *T. brichardi* and

some morphological features considered atypical for *T. sp.* ‘black’ (Van Steenberge, 2014), our new AFLP data, in concordance with Egger et al. (2007), do not find evidence for large-scale genomic admixture in these introgressed populations. There is increasing evidence that in animals local or even range wide replacement of mitochondrial DNA, without signatures of nuclear genomic admixis, seems to be more common than previously thought (e.g., Nevado et al., 2009; Tang et al., 2012; Melo-Ferreira et al., 2014; Good et al., 2015; Koblmüller et al., 2016), such that taxonomic assignment based on mitochondrial data alone might be misleading.

Acknowledgments Open access funding provided by University of Graz. We wish to thank the members of the Centre de Recherche en Hydrobiologie at Uvira, Democratic Republic of the Congo, Prof. Gaspard Banyankimbona (University of Burundi), Maarten P. M. Vanhove, Radim Blazek, as well as Meirelle Schreyen and the staff of Fishes of Burundi, for their assistance during fieldwork. This study was supported by the Austrian Science Fund (grant P22737-B09 to CS). Field work of MVS was supported by the King Leopold III Funds for Nature Exploration and Conservation who, at the time, was recipient of a scholarship of the Research Foundation—Flanders (FWO Vlaanderen). Field work of SK was supported by the Czech Science Foundation (GBP505/12/G112-ECIP).

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